Interventional treatment of venous thromboembolism: A review

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Abstract

Venous thromboembolism (VTE), including deep vein thrombosis (DVT) and pulmonary embolism (PE), is the third most common cardiovascular disease after coronary artery disease and cerebrovascular disease and is responsible for significant morbidity and mortality in the general population. Full dose anticoagulation is the standard therapy for VTE, both for the acute and the long-term phase. The latest guidelines of the American College of Chest Physicians recommend treatment with a full-dose of unfractioned heparin (UFH), low-molecular-weight-heparin (LMWH), fondaparinux, vitamin K antagonist (VKA) or thrombolysis for most patients with objectively confirmed VTE. Catheter-guided thrombolysis and thrombosuction are interventional approaches that should be used only in selected populations; interruption of the inferior vena cava (IVC) with a filter can be performed to prevent life-threatening PE in patients with VTE and contraindications to anticoagulant treatment, bleeding complications during antithrombotic treatment, or VTE recurrences despite optimal anticoagulation. In this review we summarize the currently available literature regarding interventional approaches for VTE treatment (vena cava filters, catheter-guided thrombolysis, thrombosuction) and we discuss current evidences on their efficacy and safety. Moreover, the appropriate indications for their use in daily clinical practice are reviewed.

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Introduction

Venous thromboembolism (VTE) consists of two related conditions: deep vein thrombosis (DVT) and pulmonary embolism (PE). The rate of first-time VTE events is approximately 100 persons/100,000 population each year in the United States, with this risk proportionally increasing with age, from less than 5 cases/100,000 persons below the age of 15 years to 500 cases/100,000 persons above the age of 80 years [1]. The basic mechanisms underlying VTE remain those previously described by Virchow: vascular endothelial damage, stasis of blood flow, and blood hypercoagulability; the most frequent risk factors include major surgery, trauma, hip fracture, lower extremity paralysis, previous VTE, increasing age, cardiac or respiratory failure, prolonged immobility, presence of central catheters, oestrogen treatment, and several inherited and acquired hematological conditions [2].

The latest guidelines of the American College of Chest Physicians [3] recommend treatment with a full-dose of unfractionated heparin (UFH), low-molecular-weight heparin (LMWH), fondaparinux, vitamin K antagonist (VKA) or thrombolysis for most patients with objectively confirmed VTE. However, to prevent life-threatening PE in patients with VTE and contraindications to anticoagulant treatment, such as bleeding complications during antithrombotic treatment or VTE recurrences despite optimal anticoagulation, interruption of the inferior vena cava (IVC) with a filter should be sometimes considered [3–7]. Likewise, catheter-guided thrombolysis and thromboscution are interventional approaches that may be important for the management of very selected populations.

The efficacy, safety, and the appropriate indications for these therapeutic approaches are discussed herein.

Inferior vena cava filters

Permanent IVC filters

Only two randomized clinical trials have evaluated the efficacy and safety of permanent IVC filters for the prevention of PE [8,9]. Decousus and colleagues randomly assigned 400 patients with proximal DVT and at risk for PE to receive a vena caval filter (200 patients) or not (200 patients), in addition to standard anticoagulant therapy (LMWH, UFH, VKA) [8]. The rates of recurrent VTE, death, and major bleeding were analyzed at day 12, and at two years. At day 12, two patients assigned to receive filters (1.1 percent), as compared with nine patients assigned to receive no filters (4.8 percent), had had symptomatic or asymptomatic PE (odds ratio, 0.22; 95 percent confidence interval, 0.05 to 0.90). At two years, 37 patients assigned to the filter group (20.8 percent), as compared with 21 patients assigned to the no-filter group (11.6 percent), had had recurrent DVT (odds ratio, 1.87; 95 percent confidence interval, 1.10 to 3.20). There were no significant differences in mortality or other outcomes. The authors concluded that in high-risk patients with proximal DVT the initial beneficial effect of IVC filters for the prevention of PE was counterbalanced by an excess of recurrent DVT, without any difference in mortality. The 8 years follow-up data of this population showed a rate of DVT recurrences of 34.1% in patients with filter and of 27.3% in those without (p = 0.08), while the incidence of post-thrombotic syndrome was surprisingly similar in the two groups (70.3% and 69.7%, respectively) [10].

Fullen at al enrolled in a quasi-randomised trial patients with traumatic hip fracture to receive or not a Mobin-Uddin caval filter; none of the patients received anticoagulant treatment [9]. The rate of PE was 4/41 in the filter group and 19/59 in the control group (RR 0.3, 95% CI 0.11 to 0.82), showing a statistically significant reduction in the treated group; mortality was similar for filter and no-filter patients (4/41 and 14/59, respectively; RR 0.41, 95% CI 0.15 to 1.16).

Very recently, a Cochrane review evaluated all existing published controlled clinical trials and randomised clinical trials that examined the efficacy of filters in preventing PE [11]. Only the two above mentioned studies involving a total of 529 people were included [8,9] and the authors concluded that no recommendations can be drawn. On the one hand, one study showed a reduction in PE rates, but not in mortality, and it was subject to significant biases [9]. On the other hand, the PREPIC study lacked statistical power to detect a reduction in PE over shorter and more clinically significant time periods and the trial demonstrated that permanent VCFs were associated with an increased risk of long term lower limb DVT [8].

Because of the lack of strong evidences in the literature, the latest guidelines of the American College of Chest Physicians (ACCP) recommend against the routine use of an IVC filter in addition to anticoagulants in patients with DVT; the placement of an IVC filter is suggested only for patients with acute proximal DVT or PE and a contraindication to anticoagulation because of bleeding risk [3].

Retrievable IVC filters

The vast majority of the filters which have been implanted worldwide are of the permanent type [12–15]; nevertheless, placement of such filters presents a number of long term complications. Decousus and colleagues [8] demonstrated that in high-risk patients with proximal DVT the initial benefit of IVC filters for the prevention of PE was counterbalanced by an excess rate of recurrent DVT after two years of follow-up. Moreover, one of the most important long-term complications of filters is the thrombotic occlusion of the IVC, which is reported in 6% to 30% of cases [16,17]; other significant complications include vena cava perforation, filter dislocation, migration, rupture, fracture and fragment embolization with risk of cardiac perforation and tamponade [18]. Thus, alternative strategies for IVC interruption are required, especially in patients with a long life expectancy and for whom the need for anticoagulant therapy is presumably short.

Non-permanent filters are classified as temporary or retrievable devices. Temporary filters remain attached to a wire or catheter that exits the skin; they are often difficult to manage and present frequent complications such as thrombosis, infections or migrations [17]. They must be removed within few days of placement, which is often not enough to solve the clinical problem that had led to their placement. Retrievable filters are a new generation of IVC filters and may represent a more attractive option because they may be either left in place permanently or safely retrieved after a quite long period when they become unnecessary [16,17]. This optimism must be tempered by important unresolved issues, including the appropriate maximum
implantation time, the possibility of safely and efficaciously removing the filters without being compromised by entrapped clots, and the use of anticoagulation during the implantation and peri-removal periods. A study performed in US has recently evaluated trends in the placement and removal of IVC filters in the Medicare population from 1999 through 2008 [19]. Although IVC filters were often placed not in accordance with established indications for their implantation, the frequency of their use has doubled over the past decade and the percentage of the removal was very low (ranging from 1.2% to 5.1%) [15,19]. For these reasons, the FDA has recently echoed concerns about the relative infrequency with which retrievable filters are removed and has specifically recommended that “implanting physicians and clinicians responsible for the ongoing care of patients with retrievable IVC filters consider removing the filter as soon as protection from PE is no longer needed” [20].

Different retrievable IVC filters

Results of the most important studies on retrievable filters are reported in Table 1.

Gunther Tulip filter

The Gunther Tulip filter consists of four struts of stainless wheels with hooks at the end acting as anchors. The filter can be placed either from the femoral or the jugular access, and retrieval is from the right jugular site [21]. The registry of the Canadian Interventional Radiology Association [21] and several other reports [22–26] have demonstrated the feasibility of retrieval of Gunther Tulip IVC device, after a maximum implantation time of 139 days (mean 14 days), with a low rate of recurrent PE while the filter was in place (0–3.6%); IVC thrombosis occurred in 0–9.6% and retrieval was successful in most of the cases (ranging from 88% to 98%).

ALN filter

The ALN filter is a hydrodinamic steel retrievable IVC filter. It has six short legs that ensure its adherence to the IVC walls, and three long legs that guarantee the correct central positioning into the vena cava [27]. ALN filter can be placed from the femoral, brachial or jugular vein approach, and can be retrieved only from the jugular approach. A number of trials have investigated the efficacy and safety of long-term retrieval of the ALN device, showing a maximum implantation time of 722 days (range 6–722 days) with a high rate of retrieval technical success (range 78%–100%) [27–33].

Recovery filter

The Recovery Nitinol Filter (RNF) is composed of 12 nitinol wires that extend from a nitinol sleeve and has six arms and six legs [34]. Efficacy and safety of the Recovery Filter has been evaluated in seven studies, demonstrating the feasibility of removal of the device in a high percentage of patients (retrieval technical success varying from 85% to 100%) with a maximum implantation time of 475 days (range 5–475) [35–41].

OptEase filter

The OptEase filter is a nitinol-MRI compatible filter and it is the only filter retrievable from a femoral vein approach; the filter has a symmetrical double-basket design with six straights struts connecting the proximal and distal baskets [42]. Several trials have recently investigated the efficacy and safety of retrieval of the OptEase device, showing a maximum implantation time of 48 days (range 4–48 days) with a high percentage of removal technical success (from 85% to 100%) [42–46].

Indications for filter implantation

Contraindications to anticoagulation

According to the 8th ACCP Evidence-Based Clinical Practice Guidelines on Antithrombotic and Thrombolytic Therapy [3,47], IVC filter placement is recommended when there is a contraindication or complication of anticoagulant therapy in a patient with proximal DVT or PE. Frequently, the contraindication to anticoagulation is temporary (i.e. haemorrhagic stroke, trauma) and antithrombotic therapy can be started as soon as it is resolved; for this reason, retrievable IVC filters may be the ideal “bridge” to anticoagulation for these patients. Main indications for retrievable vena cava filtration are reported in Table 2.

Table 1

<table>
<thead>
<tr>
<th>Study</th>
<th>Filter</th>
<th>Number of filters removed and placed</th>
<th>Mean duration between filter placement and retrieval (days)</th>
<th>Retrieval technical success (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Punchon, 1999 [22]</td>
<td>Gunther-Tulip</td>
<td>8 of 10</td>
<td>12; range 8–14</td>
<td>88</td>
</tr>
<tr>
<td>Offner, 2003 [23]</td>
<td>Gunther-Tulip</td>
<td>37 of 44</td>
<td>14; range 3–10</td>
<td>97</td>
</tr>
<tr>
<td>Pieri, 2003 [27]</td>
<td>ALN</td>
<td>7 of 18</td>
<td>63; range 49–192</td>
<td>100</td>
</tr>
<tr>
<td>Barrad, 2003 [28]</td>
<td>ALN</td>
<td>13 of 54</td>
<td>22; range 11–90</td>
<td>100</td>
</tr>
<tr>
<td>Pancione, 2004 [29]</td>
<td>ALN</td>
<td>28 of 96</td>
<td>72; range 30–120</td>
<td>100</td>
</tr>
<tr>
<td>Morris, 2004 [24]</td>
<td>various</td>
<td>14 of 130</td>
<td>19; range 11–41</td>
<td>93</td>
</tr>
<tr>
<td>Imberti, 2005 [30]</td>
<td>ALN</td>
<td>14 of 30</td>
<td>123; range 30–345</td>
<td>78</td>
</tr>
<tr>
<td>Grande, 2005 [35]</td>
<td>Recovery</td>
<td>14 of 107</td>
<td>150; range 0–419</td>
<td>93</td>
</tr>
<tr>
<td>Oliva, 2005 [39]</td>
<td>OptEase</td>
<td>21 of 27</td>
<td>11; range 5–14</td>
<td>100</td>
</tr>
<tr>
<td>Rosenthal, 2005 [43]</td>
<td>OptEase</td>
<td>40 of 40</td>
<td>16; range 3–48</td>
<td>100</td>
</tr>
<tr>
<td>Ray, 2006 [26]</td>
<td>various</td>
<td>80 of 197</td>
<td>19; range 1–139</td>
<td>85</td>
</tr>
<tr>
<td>Stefanidis, 2006 [37]</td>
<td>various</td>
<td>47 of 83</td>
<td>142; range 17–475</td>
<td>87</td>
</tr>
<tr>
<td>Mismetti, 2007 [31]</td>
<td>ALN</td>
<td>56 of 220</td>
<td>51; range 6–352</td>
<td>93</td>
</tr>
<tr>
<td>Karmy, 2007 [57]</td>
<td>various</td>
<td>90 of 446</td>
<td>28.2±26.3</td>
<td>78</td>
</tr>
<tr>
<td>Pancione, 2006 [32]</td>
<td>ALN</td>
<td>71 of 276</td>
<td>74; range 30–130</td>
<td>93</td>
</tr>
<tr>
<td>Pellerin, 2008 [33]</td>
<td>ALN</td>
<td>122 of 123</td>
<td>93; range 6–722</td>
<td>99</td>
</tr>
<tr>
<td>de Villiers, 2008 [38]</td>
<td>Recovery</td>
<td>22 of 54</td>
<td>48; range 7–90</td>
<td>96</td>
</tr>
<tr>
<td>Oliva, 2008 [39]</td>
<td>Recovery</td>
<td>51 of 120</td>
<td>53; range 7–242</td>
<td>100</td>
</tr>
<tr>
<td>Binkert 2009 [41]</td>
<td>Recovery</td>
<td>61 of 100</td>
<td>140; range 5–300</td>
<td>95</td>
</tr>
<tr>
<td>Onat, 2009 [45]</td>
<td>OptEase</td>
<td>124 of 228</td>
<td>11; range 4–23</td>
<td>91</td>
</tr>
<tr>
<td>Kalva, 2011 [46]</td>
<td>OptEase</td>
<td>14 of 71</td>
<td>9; range 5–21</td>
<td>85</td>
</tr>
</tbody>
</table>
Few case reports showed that retrievable IVC filters offer a safe and effective prevention to PE during pregnancy and puerperium and can be removed without complications [61,62]. On the other hand, there are no strong data supporting the routine use of IVC filters in patients suffering from acute DVT during pregnancy and this device should be reserved for selected and specific situations.

**Major surgery associated with a high risk of DVT**

Patients undergoing major orthopaedic surgery such as hip and knee replacement carry a very high risk of VTE complications [47]. Several case series showed the efficacy of IVC filters in the prevention of PE in orthopaedic patients, but none of these studies included a control group and follow-up was of limited intensity and duration [29]. Furthermore many recent advances in pharmacological prophylaxis (low-molecular weight heparin, synthetic factor Xa and thrombin inhibitors) have contributed to significantly reduce the risk of VTE in this setting. Retrievable filters remain a useful option for highly selected cases, i.e. patients at very high thromboembolic risk because of a previous, recent massive PE or recurrent VTE episodes or patients with a major contraindication to pharmacologic therapy [30]. PE is considered the leading cause of death after bariatric surgery and common pharmacologic prophylactic strategies have not been adequately tested in morbidly obese patients [50,63]. Placement of IVC filters has become a common prophylactic strategy among some bariatric surgeons, even if no prospective randomized clinical trials have compared IVC filters with alternative methods. However, filter placement can be challenging in these patients, especially in the super obese (BMI > 60). In conclusion, there are no data supporting the routine use of retrievable IVC filters in bariatric patients, and this device should be reserved for specific situations.

**Filter complications**

The most important filter complications are reported in Table 3.

**Filter occlusion and inferior vena cava thrombosis**

Occlusion of the filter is the most frequent complication of vena cava filters and its incidence varies from 6% to 30% of cases [16–19]. The reasons for this complication include thrombogenicity of the device, natural cephalic progression of DVT from the lower limb and entrapment of emboli within the filter. Thrombosis of the filter and vena cava occlusion may be associated with important clinical side effects, including decreased protection against PE, migration of the filter, post-thrombotic syndrome and chronic venous stasis. New generation filters offer the advantage of a lower thrombogenicity compared with older ones.

**Lower extremity vein thrombosis and post-thrombotic syndrome**

Vena cava filters themselves have sometimes been observed to obstruct blood flow and contribute to an increase of recurrence of DVT of the lower extremity [8,10,64]. For these reason, the ACCP guidelines recommend, if the filter is positioned as an alternative to

<table>
<thead>
<tr>
<th>Complication</th>
<th>Rate (%)</th>
</tr>
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<tbody>
<tr>
<td>Complications from insertion</td>
<td>4–11</td>
</tr>
<tr>
<td>Insertion site thrombosis</td>
<td>2–28</td>
</tr>
<tr>
<td>IVC thrombosis</td>
<td>6–30</td>
</tr>
<tr>
<td>Filter migration</td>
<td>3–69</td>
</tr>
<tr>
<td>IVC perforation</td>
<td>9–24</td>
</tr>
<tr>
<td>Post-thrombotic syndrome</td>
<td>5–70</td>
</tr>
</tbody>
</table>
anticoagulation, beginning of adequate anticoagulant therapy as soon as possible if the risk of bleeding resolves [3]. Otherwise, the optimal duration of anticoagulation in patients with permanent or optional filter that is left in situ is still uncertain. A recently published cohort study followed patients who had VTE, followed by treatment with permanent IVC filter placement and were anticoagulated long-term as soon as safety allowed [64]. Patients underwent annual physical examinations and ultrasound surveillance of the lower extremity deep veins and of the IVC filter site. Symptomatic DVT occurred in 24 of 121 patients (20%; 95% CI, 14%-28%); symptomatic PE (one fatal) was diagnosed in six patients (5%; 95% CI, 2%-10%). There were 45 episodes of filter clot in 36 patients (30%; 95% CI, 22%-38%). The rate of major bleeding (6.6%) was similar to that of a concurrent persistently anticoagulated cohort without IVC filters (5.8%). Thus, the authors suggest indefinite anticoagulation to IVC filter recipients if contraindications to anticoagulation remit. On the contrary, other data of the literature, although limited, do not seem to show significant differences in the risk of DVT recurrences after IVC filter placement with or without anticoagulation [65]. To sum up, in absence of strong evidence in the literature and waiting for the results of well-designed clinical trials, patients with IVC filter should receive anticoagulation therapy according to current guidelines in any specific clinical situation; it is not suggested to continue indefinite anticoagulation just because the filter is still present.

Finally, the association of vena cava filters with an increase of post-thrombotic syndrome is still matter of debate; the available data suggest the potential risk of post-thrombotic syndrome during long-term follow-up in patients with permanent IVC filters [8,10].

Vena cava perforation

Vena cava perforation is a usually asymptomatic complication, and without substantial clinical importance. Frequently, it is only a radiological finding which occurs when filter components extend more than 3 mm outside of the wall of the IVC [64]. More rarely, bleeding complications are associated with vena cava perforation, usually when the filter leg is withdrawn leaving an open hole; other severe consequences have been rarely reported [66,67].

Filter migration

The migration of the filter towards the heart is a potentially life-threatening complication of IVC filters, even if, in the majority of cases, migration is minor and does not result in any significant morbidity [30]. A multicenter registry found that temporary IVC filters had a dislocation rate of 4.8%; no death due to this complication was reported [58]. A recently published paper reported a high rate of strut fracture (16%) and fragments embolization (25%) of the Bard retrievable IVC filter; of interest, three out of 28 patients experienced life-threatening cardiac complications related to migration of fragments to the heart [18].

Superior vena cava filters

The placement of superior vena cava (SVC) filters to prevent PE from upper-extremity DVT (UEDVT), although controversial, has been reported. A recently published review identified a total of 21 publications that included 209 SVC filters and documented eight major filter-related complications (3.8%), including four cardiac tamponades, two aortic perforations, and one recurrent pneumothorax [68]. The in-hospital or 1-month mortality rate was 43.1%. Twenty-eight additional publications were identified which reported 3,747 cases of UEDVT. The rates of PE and associated mortality were 5.6% and 0.7%, respective-

ly. Studies imaging both upper and lower extremities found DVT 14.7 times more likely to occur in the lower extremities than in the upper extremities and the rate of PE from a lower-extremity thrombus to be 25.1%. The lack of evidence documenting the risk from UEDVT and the absence of data supporting the safety and efficacy of SVC filters bring their benefit into question.

Catheter Directed Thrombolysis

Deep vein thrombosis

Treatment of lower limb DVT should be started as soon as possible to maximally reduce the risk of further complications such as PE, recurrent DVT or post-thrombotic syndrome (PTS) [69]. PTS occurs in 20 to 50% of patients after acute DVT and leg ulceration is present in up to 10% of patients [70,71]. These conditions lead to disability and reduced quality of life, with important clinical and public health implications, as more than 50% of patients are of working age. Oral anticoagulant therapy reduces thrombus propagation, but does not effectively produce clot lysis, thus potentially resulting in an incomplete prevention of PTS, which occurs after proximal DVT in up to 50% of patients within two years [70,71]. Treatments that actively remove the clots have the potential to reduce acute symptoms and the risk of PTS by directly reversing venous obstruction and restoring the function in valves that were immobilized by the thrombus.

The effectiveness of systemic thrombolysis to achieve early clot lysis had been investigated in a number of trials which found it to be associated with high rates of serious bleeding complications with relatively modest rates of thrombus clearance [72–74]. Catheter-directed thrombolysis (CDT) involves delivery of thrombolytic agents directly through a catheter traversing the thrombus. This may be more effective in achieving local clot lysis and in restoring venous patency while significantly reducing the risk of systemic bleeding complications.

Most of the studies on catheter-directed thrombolysis are observational studies or case-series [75–82]. The long-term outcome of catheter-directed thrombolysis in patients with acute iliofemoral venous thrombosis was evaluated in 101 patients with 103 extremities affected by iliofemoral venous thrombosis [84]. At 6 years, 82% of the limbs treated with CDT had patent veins with competent valves and without any skin changes or venous claudication. In the National Venous Registry, patients with short-term thrombosis (<10 days) had better outcomes than those with older clots and correction of underlying venous lesions after successful thrombolysis, usually with intravascular stenting, appeared to be beneficial [76]. In an evaluation of 98 patients with iliofemoral DVT treated with CDT (n = 68) or anticoagulation (n = 30), quality of life was better in patients treated with CDT and correlated with the degree of lysis [81].

Few trials compared catheter-directed thrombolysis with conventional anticoagulant treatment. A single-center trial [85] randomly selected 35 patients with acute iliofemoral DVT to catheter-directed intrathrombus streptokinase or to anticoagulation alone. Six-months after treatment, patency rate was significantly higher in the thrombolysis group (72% vs 12%, p < 0.001), and the prevalence of venous reflux was lower. In a subsequent, multicenter, controlled trial, 103 patients were allocated to additional CDT (n = 50) or to standard treatment alone (n = 53) [86]. After CDT, complete lysis was achieved in 24 and partial (50%-90%) lysis in 20 patients. After 6 months, iliofemoral patency was found in 32 (64.0%) in the CDT group vs. 19 (35.8%) controls, corresponding to an absolute risk reduction (RR) of 28.2%. Although bleeding complications are the major concern with lytic therapy, recent reports have shown bleeding complication rates to be less than half the rates in earlier reports, which is likely due to more appropriate patient selection and experience with the technique. Data are not available for the comparison between different plasminogen activators or between a particular catheter or catheter-based technique to others, and there are insufficient data to assess the additional benefits of IVC filters in this setting. Overall, the results of published studies suggest that CDT may be effective in selected patients; however, indications for CDT are mainly based on...
the results of few and small RCTs. Thus, the latest ACCP guidelines suggest CDT only in selected patients with extensive acute proximal DVT (iliofemoral DVT, symptoms for <14 days, good functional status, life expectancy >1 year) who have a low risk of bleeding, if appropriate expertise and resources are available [3].

Pulmonary embolism

It has been postulated that direct, intra-embolic infusion of thrombolitics into large proximal emboli might be more beneficial than the peripheral route. In patients with acute PE, thrombolytic therapy administered directly into the pulmonary arteries does not appear to be more beneficial than that given by peripheral administration, and infusion of rt-PA directly into a pulmonary artery as opposed to a peripheral vein does not accelerate thrombolysis, but causes more frequent bleeding at the catheter insertion site. In fact, the results from a pilot trial indicate that intrapulmonary infusion of rt-PA does not offer significant benefit over the intravenous route [87]. When a lytic agent is appropriate for PE, current evidence supports that thrombolytic therapy should be infused into a peripheral vein over the intravenous route. In patients with acute PE, thrombolytic treatment should be administered via a peripheral vein rather than placing a pulmonary artery catheter [3].

Thrombosuction

Deep vein thrombosis

The combination of mechanical thrombus fragmentation (associated or not with suction) and catheter-directed thrombolysis (CDT) is frequently used in the daily clinical practice of centers performing CDT (pharmacomechanical thrombolysis). Although randomized clinical trials comparing CDT and pharmacomechanical thrombolysis are not available, retrospective data [88,89] show that the two procedures have similar efficacy and safety based on the rates of successful thrombolysis and of major bleeding, respectively. However, pharmacomechanical thrombolysis is associated with shorter treatment times, fewer days of hospitalization and is less expensive.

Based on the available literature, the latest guidelines of the ACCP suggest in patients with acute DVT the use of pharmacomechanical thrombolysis (including thrombus fragmentation and/or suction) in preference to CDT alone, with the aim to shorten treatment time if appropriate expertise and resources are available [3]. Actually, large and well-designed randomized clinical trials assessing the role of CDT and mechanical thrombectomy for reducing the incidence of post-thrombotic syndrome in patients with DVT are still ongoing (TRACT, CaVenT and DUTCH CAVA trials). However, waiting the results of these studies, pharmacomechanical thrombolysis should be used only in appropriate patients who do not have any contraindications to thrombolysis (Table 4).

Pulmonary embolism

Hemodynamically unstable PE, defined as PE with arterial hypotension or cardiogenic shock at presentation, is associated with a poor short term prognosis. Systemic thrombolysis is the mainstay of therapy in this setting, but its use remains associated with an estimated 20% risk of major hemorrhage, including a 3%–5% risk of hemorrhagic stroke. Although rarely performed, surgical embolectomy is a potential alternative treatment. However, the presence of contraindications to fibrinolytic therapy are rather common in the setting of high-risk PE, and surgical embolectomy is frequently not performed owing to excessively high surgical risk. Under those circumstances, interventional catheterization techniques (including percutaneous, catheter-based thrombectomy, thrombus fragmentation, clot pulverization, rheolytic thrombectomy, and rotational catheter embolectomy) [90–94] are reasonable alternatives for hemodynamically unstable PE. Pharmacologic thrombolysis and mechanical interventions can also be done together in absence of high bleeding risk.

There are no randomized clinical trials or prospective well-conducted cohort studies that have evaluated the efficacy and safety of interventional catheterization techniques in this clinical setting. Several case series [90–93] enrolling a small number of patients suggest that these techniques can be lifesaving in very selected clinical situations. A recently published meta-analysis evaluated the safety and efficacy of modern catheter-directed treatment (CDT) for the therapy of massive PE [95]. Modern techniques were defined, as the use of low-profile devices (< or = 10 F), mechanical fragmentation and/or aspiration of emboli including rheolytic thrombectomy, and intraclot thrombolytic injection if a local drug was infused. Clinical success was defined as stabilization of hemodynamics, resolution of hypoxia, and survival to hospital discharge. Five hundred and ninety-four patients from 35 studies (six prospective, 29 retrospective) met the criteria for inclusion. For mechanical intervention, the most common technique used was rotating pigtail fragmentation of emboli, performed in 69% of the patients (408/594). The pooled clinical success rate from CDT was 86.5% (95% confidence interval [CI]: 82.1%, 90.2%). Pooled risks of minor and major procedural complications were 7.9% (95% CI: 5.0%, 11.3%) and 2.4% (95% CI: 1.9%, 4.3%), respectively. Data on the use of systemic thrombolysis before CDT were available in 571 patients; 546 of those patients (95%) were treated with CDT as a first adjunct to heparin without previous intravenous thrombolysis. The latest guidelines of the ACCP recommend that interventional catheterization techniques should not be used for most patients with PE; this approach is reasonable only if appropriate expertise is available in selected highly compromised patients who are unable to receive thrombolytic therapy because of bleeding risk, or whose critical status does not allow sufficient time for systemic thrombolytic therapy to be effective [3].

Conflict of interest statement

All authors have no direct or indirect conflicts of interest to disclose.

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